IDEAS AND PERSPECTIVES

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Science, Policy and

Duelling timescales of host movement and disease recovery determine invasion of disease in structured populations

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Abstract

The epidemic potential of a disease is traditionally assessed using the basic reproductive number, R_0 . However, in populations with social or spatial structure a chronic disease is more likely to invade than an acute disease with the same R_0 , because it persists longer within each group and allows for more host movement between groups. Acute diseases 'perceive' a more structured host population, and it is more important to consider host population structure in analyses of these diseases. The probability of a pandemic does not arise independently from characteristics of either the host or disease, but rather from the interaction of host movement and disease recovery timescales. The R_* statistic, a group-level equivalent of R₀, is a better indicator of disease invasion in structured populations than the individual-level R_0 .

Keywords

Disease invasion, metapopulation, dispersal, epidemiological modelling.

Ecology Letters (2005) 8: 587–595

INTRODUCTION

At the turn of the 20th century, rinderpest swept through Africa, devastating populations of African buffalo and wildebeest (Sinclair 1977; Plowright 1982; Anderson 1995). From 1929 to 1983 recurrent rinderpest outbreaks occurred in the buffalo and eland populations of Central and Eastern Africa, while many other ungulate species, such as duikers, steenbok, oribi, roan, sable and gerenuk, were relatively unaffected (Anderson 1995). Why were some hosts affected more than others? Traditionally, this may have been explained by immunological differences in susceptibility. However, we illustrate a significant component of behavioural susceptibility exists that is not a simple function of group size or population density, but rather the interaction of group size and host movement.

Risk of disease is assumed to be a significant cost of group living (Freeland 1976; Moller et al. 1993), yet recent comparative analyses that investigated the effect of group size on the immune system or parasite diversity have had mixed results (Cote & Poulin 1995; Nunn et al. 2000; Nunn 2002; Stanko et al. 2002; Tella 2002; Nunn et al. 2003a,b). These mixed results may be due, in part, to the interaction of movement and group size, whereby reduced movement rates can mitigate some costs associated with larger group sizes. Specifically, large groups will be exposed to fewer introductions of disease if movement between groups is sufficiently rare.

Several recent theoretical studies have investigated the role of host population structure in the invasion or persistence of disease (Hess 1996; Swinton et al. 1998; Keeling 1999; Keeling & Gilligan 2000; Keeling & Grenfell 2000; Thrall et al. 2000; Park et al. 2001; Fulford et al. 2002; Keeling & Rohani 2002; Park et al. 2002; Hagenaars et al. 2004). These studies incorporated host movement into structured disease models either phenomenologically or mechanistically. Models with mechanistic host movement explicitly move individuals from one group to another (e.g. Hess 1996; Thrall et al. 2000; Keeling & Rohani 2002). Models with phenomenological host mixing assume that hosts do not move between groups but can infect others within and among groups simultaneously (e.g. Ball et al. 1997; Swinton et al. 1998; Hagenaars et al. 2004). The phenomenological approach may be appropriate for plantpathogen systems (e.g. Park et al. 2001, 2002), but can obscure the relationships between host movement, group size and disease recovery in mobile host populations. For example, in a system where between-group movements are rare, an epidemic of an acute, highly transmissible disease may run to completion within a group before any individual moves and spreads infection to a new group. A mechanistic model more readily captures this possibility, while a model with phenomenological mixing between groups does not. In this study, we investigate how the interactions of group size, movement and recovery affect the probability of invasion by disease into structured populations using a mechanistic mixing model.

Lloyd-Smith et al. (2004) showed that transmission of sexually transmitted diseases is well-described by a phenomenological mixing model when partner exchange is very rapid relative to the infectious period, and otherwise a mechanistic pair-formation model is required. Keeling & Rohani (2002) reached similar conclusions for a two-patch system where host mixing was frequent. We expand upon these analyses by exploring a broad range of relative timescales of movement and disease recovery as well as group and population size. Our analysis is motivated by questions regarding the invasion of disease in wildlife populations where host movement between groups can be rare or relatively frequent (e.g. natal dispersal vs. frequent fission and fusion of entire groups), infectious periods range from several days to several years (e.g. rabies vs. bovine tuberculosis), and group sizes range from monogamous pairs to thousands of individuals. We focus on directly transmitted diseases where hosts may move between groups, but contacts that are sufficient for disease transmission occur only within a group. These groups may reflect either social or spatial structure in the host population.

The basic reproductive number, R_0 , is the expected number of infections caused by a typical infectious individual in a completely susceptible population (Anderson & May 1991). The R_0 statistic has been the traditional standard by which epidemiologists and disease ecologists quantify the potential growth of a disease (Anderson & May 1991; Diekmann & Heesterbeek 2000). In stochastic models, a disease cannot invade the entire system when $R_0 \le 1$ and has a non-zero probability of invading only when $R_0 > 1$. In the simplest case of a susceptible-infectedrecovered (SIR) disease (Anderson & May 1991), Ro is the ratio of two rates, or timescales: the infection rate and the recovery rate. If transmission is density-independent, with rate parameter β , and the recovery rate γ is constant, then $R_0 = \beta/\gamma$ (McCallum et al. 2001). Further, the average length of the infectious period is $1/\gamma$. We use a stochastic metapopulation model to illustrate the importance of another ratio of two timescales, specifically the ratio of the rates at which hosts move between groups (µ) and recover from disease (γ) . For the simple case of constant recovery and no mortality, this ratio, μ/γ , is the expected

number of times an infectious individual will move between groups.

First, we describe the simulation model and explore how the interactions of group size, host movement and infectious period affect the probability of invasion by a disease. Next, we describe a relatively new metric of disease invasion, R_* , which is the number of groups that are expected to become infected from the initially infected group (Ball *et al.* 1997). In other words, R_* is the group-level analogue of R_0 . We then use the simulation model to estimate R_0 and R_* and demonstrate that R_* is a better predictor of disease invasion in structured populations with mechanistic host movement between groups. We conclude with a number of testable predictions that follow from the ideas presented here.

DEMONSTRATION OF DUELLING TIMESCALES EFFECT

We use an individual-based, stochastic, discrete-time SIR model to investigate how the duelling timescales of host movement and disease recovery affect the ability of a directly transmitted disease to invade a spatially, or socially, structured population. The total host population is evenly divided into an array of groups. The host population is further subdivided into susceptible, infected and recovered classes where S, I and R, respectively, are the number of hosts in each category. Three processes are described in the model: infection, recovery of infected hosts and movement between groups. As the intent of the model is a qualitative description of different interactions, we have simplified these processes as much as possible. For the case presented here, we consider a successful invasion to have occurred when the disease becomes a pandemic and infections occur within all groups of a structured population. This narrow definition does not count disease establishment within a single patch as an invasion, but instead emphasizes the spread of the disease among groups which is the phenomenon of interest here.

We assume that movement between groups is density-independent, and all individuals have a constant probability, μ , of leaving their current group each time step. Groups are organized on a square lattice and individuals can only move to their four nearest-neighbouring groups. To avoid boundary effects, opposite edges of the array are connected to create a torus. In Appendix S1, we expand the analysis to include a loop structure, where each group has only two nearest-neighbours, and a spatially implicit array, where individuals can move to any other group within a time step [equivalent to the 'island' model used previously (Hess 1996; Fulford *et al.* 2002)]. We assume that infected individuals recover to an immune class with a constant probability, γ , per time step.

To isolate the effects of host movement and facilitate the comparison of disease dynamics in a range of population structures, we assume disease transmission is frequencydependent (Getz & Pickering 1983). Thus the probability of infection per time step for each susceptible in group i is given by the expression

$$1 - \exp\left(-\beta \frac{I_i}{N_i}\right),\,$$

where β is the transmission coefficient, I_i is the number of infected individuals in group i, and N_i is the total number of individuals in group i. As we do not incorporate host demographic dynamics, the assumption of frequencydependent rather than density-dependent transmission represents a rescaling of the transmission coefficient β. If contact or transmission rates increase with population density (McCallum et al. 2001), then disease invasion would be even less likely when the population is divided into many small groups than indicated by results presented here, but our overall conclusions about the interaction of movement and recovery timescales would still hold.

In a continuous-time model with frequency-dependent transmission and a constant recovery rate, $R_0 = \beta/\gamma$ (Anderson & May 1991; McCallum et al. 2001). For the discrete-time model used here, β/γ is an approximation of R_0 , which works well when the probability of infection per timestep is small and group sizes are relatively large. The approximation does not change our qualitative conclusions, however, for clarity we refer to the ratio β/γ as R_0 . We also assume that disease invasion is fast relative to birth and death rates, so the total population size is constant. Each simulation starts with one infected individual, and all groups begin with the same number of individuals. As our spatial model was symmetric, group sizes remained relatively constant during the course of each run.

We begin by comparing the dynamics of two diseases with the same R_0 value (β/γ) but where one disease is slow (i.e. a chronic disease with a relatively long-infectious period; $\beta = 0.05$, $\gamma = 0.01$) while the other is an order of magnitude faster ($\beta = 0.5$, $\gamma = 0.1$). For time steps of 1 day, these parameters correspond to mean infectious periods of 100 or 10 days respectively. We simulated the invasion of these two diseases in three different host population structures: one group of 1000 individuals (equivalent to the common 'mean-field approximation' of random mixing among all individuals), 25 groups of 40 individuals and 100 groups of 10 individuals.

As expected, subdividing the population into more groups decreases the probability of pandemic (Wilson & Worcester 1945) because it decreases the average group size and increases the number of between-group jumps the disease must make to penetrate the entire population. A less obvious effect is that slower diseases are more likely to

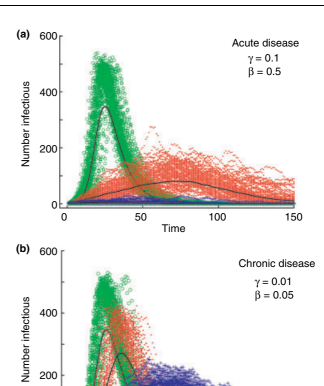


Figure 1 Disease invasion depends upon population structure (green circles: one group of 1000 individuals; red points: 25 groups of 40 individuals; blue crosses: 100 groups of 10 individuals) and the duration of the infectious period. A mean-field model of one group (green circles) is a worse approximation of a structured population for an acute disease with $\gamma = 0.1$ (a) than a chronic disease with $\gamma = 0.01$ (b). For both diseases $\beta/\gamma = 5$, but the slow disease causes more infections in the structured population. Lines represent the mean of 100 simulations. Simulations with 25 or 100 groups were run on a toroidal spatial structure with a movement probability μ of 0.01, such that $\mu/\gamma = 0.1$ (a) or 1 (b).

500

1000

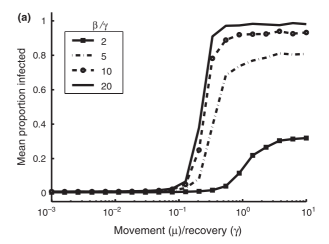
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invade a structured population, even if they have the same R_0 as a faster disease. For the case of 100 groups of 10 individuals, the slow disease ($\beta = 0.05$, $\gamma = 0.01$) infected, on average, far more individuals than the fast disease (β = 0.5, $\gamma = 0.1$) before the disease died-out (658 ± 45 SE compared with 19 ± 2.3 SE; Fig. 1). This typifies the interaction of the host movement and disease recovery timescales: diseases with longer infectious periods allow more time for host mixing to occur and thus experience populations that are effectively larger. When the movement rate is zero, neither a fast nor slow disease will invade the entire population regardless of the value of R_0 . When movement is very frequent, both the fast and the slow disease are likely to invade the structured population provided that $R_0 > 1$.

The simulated epidemics in the one-group and 100-group populations differ markedly, but were less different for the slow disease compared with the fast disease. In other words, approximating a structured population by a mean-field model (i.e. a single group with homogenous mixing of hosts) is more appropriate for slow diseases than fast diseases (Fig. 1; Cross *et al.* 2004). The faster the disease, the more important it is to incorporate the spatial/social structure into any analysis. The mean infectious period $(1/\gamma)$ defines the natural disease timescale, and when movement occurs on this timescale or slower then movement should be incorporated mechanistically, rather than phenomenologically.

Next we examine a range of host movement and recovery probabilities. The proportion of the population infected over the course of an epidemic depends on the expected number of group changes per infectious lifetime (μ/γ) , on R_0 (β/γ), and on group size (Fig. 2). When movement is infrequent relative to the recovery rate, R_0 has little predictive ability because few infections result for all values of R_0 (Fig. 2a). If movement is frequent relative to recovery, increasing R_0 increases the average proportion of the population that becomes infected (Fig. 2a), consistent with predictions from mean-field models (Anderson & May 1991; Diekmann & Heesterbeek 2000). Increasing the host group size decreases the amount of host movement required for the disease to invade the entire population (Fig. 2b). Larger groups experience larger within-group outbreaks, and hence more infected individuals dispersing from each infected group (given density-independent movement). For our model with frequency-dependent transmission, the total number of infected individuals is, on average, a fixed proportion of group size; for density-dependent transmission, the proportion infected would increase with group size, causing greater increases in the number of infected dispersers.

For high values of R_0 , the ratio μ/γ yields a sharp threshold for invasion (Fig. 2b). As a rule of thumb, a disease will invade the metapopulation if μ/γ is greater than the reciprocal of the expected number of individuals that will be infected within a single group. This makes intuitive sense because in this model system μ/γ is the expected number of between-group movements made by each infectious individual. Thus, μ/γ multiplied by the expected number of infected individuals is the expected number of infected dispersers per group, which must exceed one for a pandemic. When R_0 is high, almost all individuals in a group will be infected, so for a pandemic μ/γ should be greater than the reciprocal of the average group size. For example, if the group size is 200, then, on average, more than one infectious individual in 200 will need to move between groups (i.e. $\mu/\gamma > 0.005$) for a pandemic to occur.



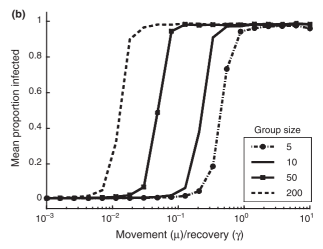


Figure 2 The interaction of movement (μ), transmission (β) and group size determines the mean proportion of the population that becomes infected. In (a) β varied from 0.2 to 2 while group size was fixed at 10. In (b) group size was increased from 5 to 200 while β was fixed at 2 ($\beta/\gamma=20$). Increasing β/γ only affected the proportion infected when movement was frequent (a). Larger group sizes require less movement for the disease to invade (b). Each parameter set was simulated 1000 times on an 11 × 11 toroidal array of groups with a constant recovery probability γ of 0.1.

The mean proportion of the population that becomes infected, shown in Fig. 2, obscures the underlying distribution of the number of infections per epidemic (i.e. over different runs of the stochastic simulation). It is incorrect to assume that the mean of this distribution is similar to the median or mode, because in many cases the distribution is bimodal with peaks centered on zero and one or close to one (Fig. 3c,f,i). When movement is very infrequent relative to recovery, the probability of a pandemic is close to zero because the disease almost always dies out within the initial group (Fig. 3a,d,g). When movement is frequent, then the disease tends to either die out stochastically within the initial

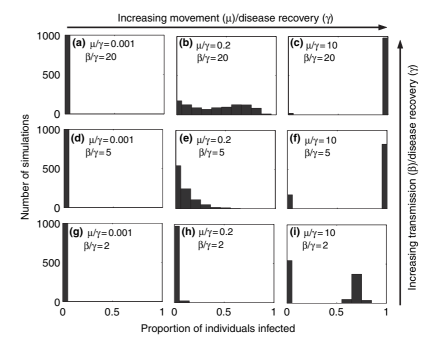


Figure 3 Histograms of the proportion of individuals infected during an epidemic for different transmission (β) and movement (μ) values scaled by the probability of disease recovery (γ). Each parameter set was simulated 1000 times on an 11 × 11 toroidal array of groups with 10 individuals each and a recovery probability γ of 0.1.

group or invade most or all of the metapopulation (Fig. 3c,f,i), with the relative frequencies of die out vs. invasion determined by R_0 as in mean-field models (Diekmann & Heesterbeek 2000). With intermediate movement rates, variation is considerable with regard to the extent to which the disease penetrates the population (Fig. 3b).

These results (Figs. 2a and 3) agree with previous studies when μ/γ is either much greater than one or close to zero. When μ/γ is large, then group structure of the population is less important and β/γ is a good predictor of disease invasion (Fig. 2a). When μ/γ is close to zero then β/γ is a good predictor of disease invasion within the initial group, but the probability of the spread of disease between groups is rather small. For the intermediate scenarios we analysed, however, the ratio of movement to recovery rate (μ/γ) has greater influence on the invasion of a disease than β/γ .

PREDICTORS OF A PANDEMIC

Recent theoretical work has extended the R_0 concept to account for depletion of the susceptible pool (Keeling & Grenfell 2000), host spatial structure (Keeling 1999; Fulford et al. 2002) and populations with heterogeneous infectiousness or susceptibility (Diekmann & Heesterbeek 2000). However, even after incorporating these effects R_0 may be misleading in metapopulations with limited mixing because, R_0 as is traditionally used, is an individual-based measure. Ball et al. (1997, 2004), Ball (1999), Ball & Lyne (2001) and Ball & Neal (2002) demonstrated that the individual-based R_0 is not the best predictor of disease invasion in structured host populations and introduced a group-level reproductive number, R_* , which is the average number of groups infected by the initially infected group. This finding has been echoed in the context of reproductive fitness of a new mutant in a metapopulation (Gyllenberg & Metz 2001; Metz & Gyllenberg 2001). In a model with phenomenological mixing, $R_* > 1$ is the formal threshold criterion for invasion of a disease into an infinite number of finite-sized groups (Ball et al. 1997).

The phenomenological mixing model used by Ball and colleagues facilitates analysis, but to demonstrate the utility of the R* metric in the context of interacting timescales of host movement and recovery, we applied our simulation model with explicit host movement between groups. For the model described above, we estimated R_0 and R_* by tracking the mean number of infections caused by the initially infected individual or group respectively. Then we averaged these estimates of R₀ and R_{*} over many runs of the stochastic model. When a susceptible individual was infected and two or more infected individuals were present within the group, we randomly allocated the infection to only one of those infectious individuals. To estimate R_* , we tracked the number of groups that were infected by individuals that were themselves infected within the index group. Infected individuals had to move to a completely susceptible group and cause infection there in order to contribute to R*. When individuals from multiple groups moved to a susceptible group and caused an infection, we randomly allocated the infection to one of the individuals (and thus its source group). The mean estimates over many simulations, denoted as \hat{R}_0 and \hat{R}_* , are 'empirical' in the sense that they are based on data collected from simulated epidemics mimicking epidemiological contact-tracing data. As estimates from model output, they incorporate the effects of spatial structure, host movement, and depletion of the susceptible pool. Thus they will differ from traditional analytical R_0 and R_* values, which assume an infinite susceptible population and, hence, overestimate the value of these parameters when populations have a finite size.

We simulated the model using a range of transmission (β) and movement (μ) probabilities, a fixed recovery probability $\gamma = 0.1$, and an 11 × 11 toroidal array with 10 individuals per group and nearest neighbour movement. Each parameter set was simulated 1000 times to generate mean values of \hat{R}_0 and \hat{R}_* . We then plotted the relationship between the model output variables \hat{R}_0 , \hat{R}_* , and the proportion of the population infected (Figs 4 and S2). Each line in Fig. 4 corresponds to a

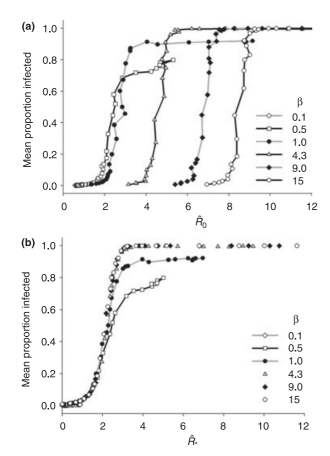


Figure 4 \hat{R}_0 can be substantially greater than one and yet not cause a pandemic (a), whereas \hat{R}_* is a much better single predictor of the mean proportion of individuals infected (b). Each line represents a fixed transmission parameter β and a range of movement probabilities from zero to one (increasing from left to right) sampled on a log scale. Each parameter set was simulated 1000 times on an 11×11 toroidal array of groups with 10 individuals each and a recovery probability γ of 0.1.

fixed within-group transmission rate (β) and a range of movement probabilities increasing from left to right.

The empirical individual-level \hat{R}_0 is not a good predictor of the mean proportion infected: even when R_0 is much greater than one, the mean proportion infected may be low depending upon the movement probability (Figs 4a and S2). Also for different β , the proportion infected appears to show a threshold at different values of \hat{R}_0 . The group-level \hat{R}_* , on the contrary, is a much better predictor of a pandemic in a structured population (Fig. 4b). In an idealized metapopulation, $R_* > 1$ is the threshold above which there is a finite probability of disease invasion (Ball et al. 1997). In our simulations, the proportion infected begins climbing at $\hat{R}_* \approx 1$ and rises most steeply around $\hat{R}_* \approx 2$ (Fig. 4b). This gradual transition around the threshold is typical of stochastic epidemic models, particularly with spatially constrained mixing, because the invasion has many chances to die out before invading the entire population. When transmission rates (and hence R_0) are low, \hat{R}_* is small for all values of movement (Fig. 4b, $\beta = 0.1$). When transmission is intermediate and movement is frequent, the disease will either stochastically die out in the initial group or invade the entire population (Fig. 3d-f), resulting in intermediate values of \hat{R}_* and mean proportion infected (Fig. 4b, $\beta = 0.5$ and 1). Finally, when both movement and transmission rates are high, \hat{R}_* and the mean proportion infected are also high.

FUTURE EMPIRICAL RESEARCH

These findings suggest important directions for empirical studies, as well as a number of testable predictions. Previous analyses of disease presence/absence in different host social structures have not considered the interaction between movement rates and the duration of the infectious period (e.g. Cote & Poulin 1995; Nunn et al. 2000; Altizer et al. 2003; Nunn et al. 2003a,b). However, our results illustrate that it is the relative timescales of movement, recovery and infection that determine the probability of a pandemic. A slow, chronic disease may 'perceive' a host to be relatively well-mixed with frequent movement of individuals among groups. An acute disease will perceive that same host population to be more structured because movements between groups are less frequent relative to the timescale of the infectious period (Cross et al. 2004). We hypothesize that all else being equal, chronic diseases will be more likely to penetrate structured populations than acute diseases. Conversely, we hypothesize that behaviourally susceptible host species, with large groups and frequent movement, are likely to be more heavily impacted by acute diseases than hosts with small groups and infrequent movement. Thus the ratio of acute to chronic diseases found in different host populations should increase as a function of group size and movement rate.

A major focus of recent disease ecology has been how transmission or contact rates depend on population density (e.g. Bouma et al. 1995; Begon et al. 2003), but for metapopulations, we have shown that movement rates are also critical to understanding disease invasion (Figs 2-4). Despite the importance of host movement, very few studies have been published that examine the amount of mixing between groups of many wildlife species. Our results can help to guide the design of field studies intended to estimate host movement for disease models. The proportion of individuals that should be tracked and the duration of the study will depend upon the infectious period of the disease as well as the average group size of the host. As group sizes and infectious periods increase, the amount of movement required for a pandemic to occur decreases. Low movement rates, however, will require researchers to track more individuals to accurately estimate the amount of movement between groups. If group sizes are large, for example 200, and the disease is highly infectious, for example $R_0 > 5$, then only one in c. 200 infectious individuals needs to switch between groups for a pandemic to become likely (Fig. 2b).

Researchers may estimate movement between groups using genetic data or tracking of known individuals (Waser et al. 1994; Koenig et al. 1996; Cain et al. 2003; Nathan et al. 2003). Radio-tracking or resighting data are more effective than genetic methods as long as individuals frequently move between groups relative to the duration of the study. Genetic methods of estimating movement will be relevant only for chronic diseases in large groups, and their use in a disease context involves at least three major assumptions: (i) past movement accurately reflects current movement; (ii) short-term movements that are likely to be missed in genetic signatures (e.g. foraging rather than mating) are unimportant to disease dynamics; and (iii) moving individuals are as reproductively successful as non-moving individuals (Waser et al. 1994).

FUTURE THEORETICAL RESEARCH

Our findings emphasize that the group-level reproductive number R* is a critical determinant of invasion success in structured populations. Analytical formulations of R* in systems with explicit host movement may clarify the important interaction between timescales of host movement and disease recovery, and help to formalize the rule of thumb proposed above. Previous work on R* has focused on models with phenomenological host mixing and an infinite number of groups such that all new infections are in susceptible groups (Ball et al. 1997, 2004; Ball & Neal 2002). Analogous to recent developments in the theory of R_0 (Keeling 1999; Keeling & Grenfell 2000), further work on R* should consider finite populations with spatial constraints on movement. Longer dispersal distances and spatial configurations that increase the number of neighbouring groups (Appendix S1) will mitigate the depletion of susceptible groups and facilitate the invasion of a disease. These effects are implicitly incorporated into our R_0 and R_* estimates, but analytical exposition would help advance our understanding.

In our stochastic model, we made a number of simplifying assumptions that could be relaxed to make our simulations more realistic. First, the assumption of a constant probability of recovery per time step, γ, results in a geometrically distributed infectious period. The effects of alternative infectious period distributions on R* are unclear (see Keeling & Grenfell 2000). For instance, with a fixed infectious period, time spent in the home group while infectious will increase the number of local infections, but will also diminish the infectious period in any new group, thereby decreasing the number of infections elsewhere. A fixed infectious period would also cause fewer individuals to recover before moving, compared with the geometric infectious period (with its mode at one timestep). Second, we have assumed that movement between groups occurs instantaneously and without mortality, but if individuals spend time or die during movement their infectious lifetime within the next group is reduced, thereby decreasing R*. Finally, we assumed that disease invasion was fast relative to the timescale of host birth and death. This is less likely to hold for chronic diseases, or for acute diseases that lead to rapid mortality rather than recovery. Both natural and disease-induced mortality shorten the infectious period and thus reduce R_0 (Anderson & May 1991) and R*. Our broad conclusions about the interaction of host movement and disease recovery timescales should still apply, but investigating the effects of host demographics and disease mortality on R* would be an important extension of this study.

CONCLUSION

Traditionally, epidemiologists and disease ecologists have focused on $R_0 > 1$ as a threshold for disease invasion (e.g. Anderson & May 1991; Diekmann & Heesterbeek 2000). We have shown that in metapopulations the relationship between invasion of disease and an individual-level R_0 is often weak. Even for very large values of R_0 , a pandemic is unlikely if the expected number of times an individual will move between groups during their infectious lifetime (μ/γ) is low (Fig. 2). Pandemics in structured populations require both within-group and between-group transmission, and the group-level reproductive number R* is a better predictor than the individual-level R_0 for these systems (Fig. 4). Results from our individually based stochastic model support the analytical results of Ball et al. (1997, 2004) and Ball & Neal (2002), which proved that $R_* > 1$ is the threshold for disease invasion in a population with group structure. As a general rule of thumb, the individual-level R_0 must be >1 and the expected number of group changes while infectious (μ/γ) multiplied by the average group size must be >1 for a pandemic to occur (Fig. 2b).

Chronic diseases with longer infectious periods allow for more mixing to occur between groups. As a result, chronic diseases will perceive more thoroughly mixed host populations and exhibit dynamics that are closer to those predicted by mean-field models than acute diseases (Fig. 1). For the same R_0 , chronic diseases are more likely to invade structured populations than slow diseases. 'Slow' and 'fast' diseases are relative terms: a fast, acute disease in a host population with frequent movement between groups may behave like a relatively slow disease in a population with less frequent movement. The probability of a pandemic in a structured population is thus an emergent property of the interaction of host and parasite demography and behaviour, incorporating a dimension of host behavioural susceptibility arising from group size and movement rates. The results presented here, and in a recent paper by Lloyd-Smith et al. (2004), suggest that when contact, movement, birth and death processes occur on a timescale similar to that of the disease (i.e. the infectious period), these processes should be incorporated mechanistically into disease models.

ACKNOWLEDGEMENTS

This research was funded by the NSF-NIH Ecology of Infectious Disease Grant DEB-0090323 (PCC and WMG), NIH-NIDA Grant R01-DA10135 (JLS and WMG), and a James S. McDonnell Foundation 21st Century Science Initiative Award (WMG). Previous versions of this manuscript were greatly improved by the comments of Wendy Turner, Shirli Bar-David, George Wittemyer, Charles Nunn, Molly Smith and three anonymous referees.

SUPPLEMENTARY MATERIAL

The following material is available from http://www.blackwellpublishing.com/products/journals/suppmat/ELE/ELE760/ELE760sm.htm

Appendix S1 Extension of analysis.

Figure S1 Effect of population size and configuration.

Figure S2 3-D plot of \hat{R}_0 , \hat{R}_* and proportion infected.

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Editor, Jonathan Sherratt Manuscript received 22 December 2004 First decision made 24 January 2005 Manuscript accepted 9 March 2005